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Hemarks:

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- (54) Polymerisation catalyst
- A polymerisation catalyst comprising:

(A) rare earth metal component selected from the group consisting of (1) a reaction mixture obtainsble by treating a silica support, a carbon black support or a mixed support thereof with a mono- or diethyl aluminium chloride, adding a necdymium carboxylate in the presence of a solvent to form a sturry, and removing the solvent or (2) a reaction mixture obtainable by dissolving a neodymlum carboxylate in a first solvent, adding silica or carbon black, and removing the solvent to obtain a dry solid, contacting the dry solid with a solution confaining a mono- or di-ethyl aluminium chloride in a second selvent, and removing the second selvent:

(B) a co-celelyst selected from the group consisting of a disobutyl aluminium hydride, trilsobutyl aluminlum, and a mixture thereof.

Description

Field of the Invention

This invention relates to a process for producing polydienes in a gas phase reactor. More particularly, the invention of the particular of the process of a rare certificated stativet.

Background of the invention

Polydianaes such as polybyladiction a and polysisprene have been manufactured for many years by exhibitor polymorticalism and more recently by mass or bulk polymorization processoe. Various existylic activition and butk or mass processes for this polymorization of beteriam are known in the art to be suitable for producting polybuladiens with a right of polymorization of the polymorization of beteriam are known in the art to be suitable or producting polybuladiens with a right of polymorization production and polymorization are polymorized to the manufacture of tiles, betting, and other noticed or extruded to higher or electronic and one of the polymorization and polymorization and polymorization and polymorization.

In a skulion polymerination behanken in polymorized in an inert andvert or dikteral which does not enter into the struct are of or advantage intelled, the researching polymer. Such adventure nor usually disphair, command and sydonal/pinkel furytiocations section is pentame, horame, heptame, benzene, fotener, cyclohexarse and the IRo. In bolk polymerizations, the recitotion medium is essentially exhantenes, and the memorane's comployed as a dilutent.

The discovery of gas-phises faildized bed and stirred reactor processes for the production of polyment, especially polyatelian polymens, made it possible to product a wide variety of new polymens with highly declarable and improved properties. These papers provided a majors for producing such polymens provided a majors for producing polymers with a drastic production in explain investment expense and deshable deshable of the programment of the production of

as in a conventioning pas fluidisced bed process a generous serieums containing on or more monomers is passed into intellized bed treated recreating a held of growing polymer particles in applywrizedation zone, while confirmantly of non-mitternily including a polymerization content in the confirmation of the polymerization and in proceeding and the polymerization recreation is excellent than the polymerization recreation is excellent the excellent than the polymerization recreation is excellent than otherwise, distributional than the polymerization recreation is excellent than otherwise, distributional than the polymerization zone which much be removed to prevent the polymerization cone which much be removed to prevent the polymerization to provide intelligent and study applicability. This is accomplished by continuously removing unremarked that gives entitle to continuously and the polymerization zone where the polymerization zone was compressed, cooled in a head evoluting, as applicational by additional amounts of minimate the propose mornouser polymerization and manufaction that recording of the recording of

Conventioned gas phase fluidized bed resin production is very well known in the ent as shown, for example, by the discourse appending in United States Patent Nos. 4,379,766; 4,383,085 and 4,876,380, which are incorporated herein by reference.

The production of polymeric substances in gas phase stirred reactors is also well known in the art as exemplified by the process and equipment descriptions appearing in United States Patent No. 3,258,263.

More rewently, In U.S. Petient Nosc., 4,986,564 and 1,5,046,568. It has been taught that stelly polymers, including polybatterians publisher, can be produced in a fluid road but respect in this presence of a catalyst in a polymers function solver the solitering temperatures of the steller polymers in the presence of an inet particulate material. The stildy polymer produced in the gas phase process are granular having a mixture of subset and land material. The stildy polymer produced in the gas phase process we may be applied of their instellar. Purifice, U.S. Patent No. 6,377,058 discloses gas phase polymerization processes which utilize unsupported, edubtio catalysts, such as, for exempting materials conclination catalysts. The catalysts are introduced into the reaction, such as a falliable doed, on a columnia of their produced by the process may contain disease. Po 647,657 A1 discloses supported rare earth catalysts for as shore polymerization of confusion of their columnia of the process of the state of the process of the

For many years it was erronously believed that to allow tipul of any first to enter his the polymerization region of a ges piece reactor would invalidely lead to applicate about of each purisher, formation of large polymer churchs and utilizately complete reactor that down. This consens caused gas phase polymer produces to carefully exclod cooling the proyed gas stream antering the reactor to a temperature halow the condensation temperature of any of the inconcers conscious of the advanced not reaction.

Commonates such as however, i. 4-millely-partition, and oclaims 1, are partitionally valuable for producting deligent outpolyment. These higher alpha claims here relatively high condensation the imprendence. Due to the upperhandon that ligited uncommes in the polymerization zone would load to aggiomeration, churcking and ultimately shut claver the readtor, production rates, which depend upon the rate of a virth heet is removed from the polymerization zone, were asserted constrained by the perceived need to maintain thus temperature of the cycle gas stream entanting the reactor of temperature and the production of the cycle gas stream in the cycle gas stream.

Even in the case of polymerization reactions conducted in attired reactors, care was exercised to maintain the resin bed temperature above the condensation temperature of the recycle gas stream components.

To maximize heat reviewed it was not unusual to group or leight Equid folior or not the polymer had where it would immediately tach into a greeous state by exposure to the hotter recycle gas stream. A finited amount of admittental cooling was exhibited by this technique by the Jourse-Thompson offset but without ever cooling the recycle gas stream to a level where condensation might occur. This approach typically involved fine federicus and energy wasting approach of peganetry cooling a portion of the cycle gas stream to obtain liquid monomer for intergral and eutosequent separate introduction into or onto the polymerization bed. Examples of this procedure are found in United States Patent Nos. 3,294-0/07.8 (300-475) 365-285.7 and 4,012-678.

It was discovered later, contrary to the long held belief that the presence of liquid in the cycle gas stream would lead to agglomeration and reactor shut-down, that it is indeed possible to cool the online cycle gas stream to a temperature where condensation of significant amounts of monomer would occur without the expected dire results when these liqulds were introduced into the reactor in temperature equilibrium with the recycle gas stream. Cooling the entire cycle gas stream produces a two-phase gas-liquid mixture in temperature equilibrium with each other so that the liquid contained in the gas stream does not immediately tach into vegor. Instead a substantially greater amount of cooling takes place because the total mass of both gas and ikipid enters the polymerization zone at a substantially lower temperature than previously thought possible. This process led to substantial improvements in the yield of polymers produced in the gas phase, especially where compromers which condense at relatively low temperatures are used. This procedure, commonly referred to as "condensing mode" operation, is described in detail in United States Patent Nos. 4,543,399 and 4,588,790 which are incorporated by reference. In condensing mode operation the two-phase gas liquid mixture entering the polymerization zone is heated quite rapidly and is completely vaporized within very short distance after entry into the polymerization zone. Even in the largest commercial reactors, all liquid has been vaporized and the temperature of the then totally gaseous cycle gas stream rated substantially by the exoffrence nature of the polymeriza-If on reaction soon after entry into the polymerization zone. The ability to operate a gas phase reactor in condensing mode was believed possible due to the rapid healing of the two-phase gas liquid stream entering the reactor coupled with officient constant back mixing of the fluidized bad leaving no liquid present in the polymer bed more than a short distance above the entry level of the two-phase gas-liquid recycle stream.

We have now found that liquid monomer may be present throughout the entire polymer bed provided that the liquid monomer present in the bed is adsorbed on or absorbed in solid particulate matter present in the bed, such as the polymer being produced or fluidization aids present in the bed, so long as there is no substantial amount of free liquid monomer. This discovery makes it possible to produce polymers in a gas phase reactor with the use of monomers having continuation temperatures much higher than the temperatures at which conventional polyoletins are produced in gas phase reactors. Another way of viewing this discovery is that it is now possible to produce polymers using readily condensable monomers (e.g., 1,3 butadiene, having a normal boiling point of -4.5°C) in a gas phase reactor under conditions at which the monomer would be expected to be present as a liquid. Furthermore, it had been previously believed that gas phase processes for producing polymers with some or all of the monomers having low to moderate condensation temperatures were impractical because the amount of polymer produced per catalyst particle was too low at all monomer concentrations that had condensation temperatures below the temperature in the polymerization zone. The discovery of this invention now makes it economically practical to produce polymer with monomera at concentrations where they frave condensation temperatures higher than the temperature in the polymerization zone, such that liquid monomor is present throughout the entire polymer bed provided that the liquid monomer present in the bed is adsorbed on or absorbed in solid particulate matter, the polymer bed, and/or the forming polymer product present in the polymer arization zone of the reactor. This invention makes possible the gas phase production of classes of polymers which pre-50 Viously were inqualit not capable of production in a continuous gas phase process.

Another benefit of the financian is that operation with monomer present as liquid dissolved in the polymer gives a greater concentration of monomer at the nective catigated to their operation with monomer on dissolved. Let, present only in the gas please. This should modulate the productivity of the catalyst for making polymer. Still another benealt of the invention is this heat transfer within the polymer perificise should be improved use for removal of heat by monomor see evaporation. This should lead to more uniform polymer protective, produces, more uniform polymer greateries, produces, more uniform polymer protective, first produces of the processity improved polymer morphology them operation with recommend not dissolved, i.e., present only in the gas please.

Summary of the Invention

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The present invention provides a process for producing polybutatilens or polylooprene in a stirred bed or gas fluidized polymerization vessel having a polymerization zone under polymerization reaction conditions, which process comprises:

- (f) introducing buladiene or isoprene monomer into said polymerization zone containing a bed of growing polymer particles in the presence of an inert particulate majorial and collonally at least one inert gas;
- (ii) continuously or intermittently introducing a polymerization catalyst containing a rare earth metal component, a co-catalyst, and optionally a promoter into said polymerization zone;
- (III) continuously or intermittently withdrawing polybuladiene or polyscoprene product from said polymerization zone; and
- (iv) withkrawing urreacted butations or isoprens from said polymerization zone, compressing and cooling said butations or isoprens and said inert gas when present, while maintaining the temperature within said polymerization zone below the dew point of the monomer present in said polymerization zone.

Granular particles and articles produced using such particles are also provided.

Brief Description of the Drawing

A fluidized bed reaction system which is particularly suffect to the production of polybutadiene and polyisoprene is illustrated in the drawing.

Detailed Description Of The Invention

While not limited to any particular type or kind of polymerization reaction, this invention is particularly well saided to object the conclusions involving benopolymerization and expolymerization of relatively high boiling or readily condensation moments such as butdefluen and feoppere.

Examples of higher boiling or readily condensable monomers depable of undergoing definite polymerization readtions are the following:

A. higher molecular weight alpha olelins such as decene-1, dodecene-1, isobutylone, styrene and the like.

 B. dienes such as hexadiene, vinyl cyclohexene, dicyclopentadiene, butadiene, Isoprene, ethylidene norbornene and the fike.

 C. poter vinyl monomers such as acrylonitrite, maleic acid esters, vinyl acetate, acrylate esters, methacrylate esters, vinyl trialityl silanes and the like,

These higher builing or readily condenselba monomers can be homopolymerized in accordance with this invanion with this man of an intent gan as the graceries correposed not be two phase particular division cognition than the reador. 40 Publish intent majoritat for this purposa include nitrogon, argin, not caturated hydrocarbons which remain gaseous at a temperature and below the temperature selected to be metaltated in the purpoyrectation zone.

The higher holling or resulty continensitie monomers can also be copolymerized with one or more lower boiling monomers cuch as orbytone, propylene and bottone, as well as other higher boiling monomers exch as those mentioned above, the only requirement being that these be a suitilisent difference in the confinential retrigionations of the higher boiling or resulty confidentiable monomer and at least one lower boiling monomer or inet substance as will ellow enough gus to be precent in the veyel age set expent to persing legantials, elseey detail, continuous operation.

In accordance with our invention the higher boiling or readily condensable noncomers can be directly intendiated in the polymerization zone or windful lated the polymerization zone or with the ecycle gas extend or a combination of both. In preferred embodiment the temperature within suid-polymerization zone is ministanced below the condensation of both. In preferred embodiment the temperature within suid-polymerization zone is ministanced below the condensation of the condensation and the properation of the preferred embodiment the conditions (e.g., emperature, pressure, monomer(s) concentration) within said polymerization zone are used that occupilations (e.g., emperature, pressure, monomer(s) concentration) within said polymerization zone that has necessation zone are maintained such that a portion of the monomer in the province that the province zone are maintained such that a portion of the monomer in the province that the province zone are maintained such that a portion of the monomer in the province that the province zone are maintained such that a portion of the monomer in the province that the province zone are maintained such that a portion of the monomer in the province zone are maintained such that a portion of the monomer in the province zone are maintained as the provinc

The catalyst supplyed in the polymericalize zone is a ware set in made catalyst. The precise of this invention has instituted on any precised values of one use with mead of only probled that have been proviously employed in surry, solution, or bulk polymerications of higher boiling or readily contensable monomers (e.g., busideline end has provide one of the provide provid

a metal component, a co-catalyst, and optionally a promoter. Perfernitly, a promoter is not employed in the entalyst of its polymerization process of this invention. The metals component can be a nere earth compound on a militarie of two or more aree earth metal compounds, in general, the race earth metal component of the catalyst can be could be not inconcision, and the catalyst can be could be not inconcision. The catalyst can be considered or unsupported, or unsupported, or only defed in either this presence or absence of a filter. Afternatively the catalyst can be introduced to the polymerization zone in the term of a prepalymer using techniques shown to those saltied in the

When the metal component is engineted, typical supports an include, for consentate, callica, carbon black, portune crossificial polystycen, protous crossificial protous cross

Any compound, capanic or inorganic, of a motel chosen from those of Circup IIIB of the Periodic System Rendrig and elember terminal or between 57 and 100 can be employed herein. Exemples of rise earth invelae compounds are compounds of cockens. Enriforment, presecdymium, geololisium and necepturium. Of these compounds, carboxylettes, calorise, activities, calorise conspicuates, haides (including altern and nelsool completes on incondument including), and willy deliverables of the motels are preferred. Neceptum compounds are the most preferred. Neceptum compounds can include neceptum registries in neceptum in a conductiva neceptum in a compound in tribitation of the compounds of the conductivate of

The eatalyst modifiers and co-catalysts consist of aluminum alkyl halides and trialityl aluminum compounds such as the following:

Alkylatuminum halides can be a compound having the formula $AH_{(0,0)}X_n$ wherein each F is independently alkyl having 1 to 14 carbon atoms, each X is independently chlorine, promitine, or faciline, and g is 1 or 2 or a mixture of compounds having the formulae $AH_{(0,0)}X_n$, and $AH_{(0,0)}X_n$ interes in X, wherein F, X, and AH are the same as above.

Examples of helogen contining modifiers and counterfue are distriptional modifieds, ethylataminum sescychloride; cir- budyalminum miloride; dilaboutylatunium orbioride; metrylatunimum escrycthoride; isobudyalminum num escrychloride; cimetrylatunium chloride; cii-epropylatunium orbioride; metrylatunimum ciloride; ond lobobuglatunium dichloride. Dietylatunium chloride (DEAC) and disobutylatunium chloride (DIAC) are most pretered.

The fieldefuluminums can be a hydrocarby as follows: Irisolophyllouninum; thhosphluminum chickolophyllouphinum, licholophyllouphinum, licholophyllouphinum, licholophyllouphinum; licholophyllouphinum; histolophyllouphinum; birtsolytikuminum; birtsolytikuminum;

Prefereç Lo catelycht fint can be enjolyeyd with rare en'in moial compounds inclured trially alluminum (TEAL), the loobitylatuminum (TEAL), thinkeylatuminum (TEAL), the preference (MAC), modific methylatuminum (MACA). 46 threathylatuminum (TMA), a dialkyl etuminum hydride or a nature of a dialkyl etuminum hydride end a Iralkyl etuminum.

When MAG or MAMC is employed as the co-catolyst. It may be one of the following: (a) branchest or cycle objection more polythydrocauthylatuminum and total written contain respecting units of the general formatic ACM(ET)Or, where PT is hydrogen, an ally inaction aromatic action to the containing from 1 to about 12 subons atoms, or an any matical certain as a substituted or unstained principle or reportibly group; (b) bring subsitisf the general formatia fa/*IgBH*2,) where AT is a callionist bewise of Branched add enginested or obstacting an ality, inleapen, or hydrogen from the metal component of the catalyse, B branched and component of the catalyse, B branched and component when AT is a substituted convexie bydrocarbox, prelambly a perferencement product at (a) boron allysis of the general formatia BTA*, where PT is as edited add one, and an additional production and the production of the prelamble of the general formatia BTA*, where PT is as edited above.

Aluminoxanes are well known in the art and comprise oligometic threat alleyt atuminoxanes represented by the forfact india. 39

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10 and oligomeric cyclic alkyl aluminoxanes of the formula:

20 whoods §E 1 fr. 40, preferably 10 to 20; p is \$1 + 40, preferably 10 to 20; and IP* is an ellyl group containing 1 fo 12 carbon norms, proferably methyl or an avyl ediscion both no an abellited or unsubstituted ellerphyl or neightful allocal. ModRied methylatuminssom is formed by abellituding 20-00 wife of this multilyl groups with n C₂ to C₁₂ group, preferably with toolulityl groups, using enchangues known for lone soldied in the sign.

Promoters that can be used with rare earth metal compounds include Lewis acids such as BCl₃, AlCl₆, althylatuminum dichloride, althylatumis sequichborde, diethylatuminum chloride, and other alkyl radical derivatives of these compounds. Organohalide derivatives of the formula

in which X is Cl or Br, Ris H, elkyl, aryl, alkylaryl, distore or bronne alkyl, alkoxy or epoxy; R' is elkyl, aryl. H, Cl or Br; R' is elkyl, aryl. H, Cl or Br; B' is elkyl aryl. H, Cl or Br; B' is elkyl. H, Cl or Br; B'

Typical examples of organishalide derivatives which can be used as catalytic components of the present invention are bearbyl, propley), beavyl, beavyl, beavylidene or tertilary buyly chlorides or bromides, methyl chloroformate or bromoternate, chlorofitybenymethane or chlorofichenymethane, and the life.

The catalyst cem be prepared by mishing this support material, the metal component, containyst, epitonal promoter in any order in a mind relevance of tilensh, in granted the metal component can be interpretated or a support of the second of

A preferred method for making this catalyst of this investion broduce impregnating a sides support, a notifical between perfort or included support of the New with a zero early medic continued to making incomposed. This amount of motal impregnated on this support can incure between 0.1 and 1.0 motivity analysts, an organize aligit aluminum conspound to endown term is at IRF. The catalyst may be foolisted as a dry selfer or used no a clutter in a hydrocarbon or evogeneeted confront certain as IRF. The catalyst may be foolisted as a dry selfer or used no a clutter in a discoverance of the most white of the catalyst may be foolisted or the confront certain as the catalyst may be foolisted the control of the confront certain and the confront certain certain confront of the most self-time or entry which is fell discovered to the reactor. The A1 or metal may be used into discoverance of the confront certain cer

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the reactor with the cocatelyst. When MAO is used as the occatelyst no halide source is required.

A preferred procedure for making the rare earth catalyst of the invention comprises the sequential steps of (A) treating a silica support, carbon black support, or mixture of the two support materials with a mono- or di-elliyi aluminum. chloride or a mixture of the two chlorides in a hydrocarbon solvent thereby forming a slurry; (8) adding the rare earth compound (e.g., a needymtum compound); and (C) removing the hydrocarbon solvent or diluent. Catalysts which are preferred in the process of this invention are (i) a needymlum needecancate, needymlum octoate, or needymlum versatate as the metal component, and an organic alkyl aluminum compound such as diethyl aluminum chloride to form the catalyst in a diluent such as n-hexane or cyclohexane and (II) a nectlymlum carboxylate or alcoholate such as neodymium needegenoate, needymium octoale, or needymium versalate as the metal component is impregnated on sitical by dissolving the needymlum compound in THF, adding silice, followed by solvent removal. The dry solid is added to a hydrocarbon (e.g. hexane) solution containing an alkylatuminum chloride with subsequent removal of the hydrocarbon solvent. These catalysis are ted to the reactor with a co-catalyst selected from the group consisting of discoulyl attiminum hydride (DIBAH), triisobulylaluminum, or a mixture of diisobulyl eluminum hydride and trusobulylaluminum (TIBA). These catalysts are preferred because they have little or no induction period and remain catalytically active for a long period of time. The catalyst I above can be fed directly to the reactor. Still another cutalyst can be prepared by forming a reaction mixture by (I) contacting a necelymium compound selected from the group consisting of a needymium carboxylate, a needymlum alcoholate and a recdymlum acetylacetonate with a mono ethyl aluminum dichloride, a di-ethyl aluminum chloride or a mixture of the mono- and di-clin) aluminum chloride (ii) depositing the mixture on a silica support in the presence of a solvent to form a sturry aix! (iii) removing sald solvent; and adding a co-catalyst selected from 20 the group consisting of (I) dialkyl aluminum hydride, (ii) a trialkyl aluminum, (III) a mixture of a dialkyl aluminum hydride and a triality) aluminum, (iv) methylaluminoxane, (v) modified methylaluminoxane, (vi) and mixtures thereof.

and a party mannaum, (a) menylauramount, (c) mounter menylauramount, (c) and manufacture treat.

Flidiciation has employed in the breation can be daring periodule menterals which are adminishly herit for the reaction. Examples of such fluidication aids include earlion beek, ellips, days and other file meterials such as talls. Organic
polymeric menterials can also be veryloped as a fluidication aid. Caston blacks and ellibar are the professer distribution aids with cathon blackship the most professer. The earlien black materials employed have a primary purifice daze of
both 110 to 100 monovibes end an exercise size of argregate forinary enforcintery of proton. (1) to about 10 micross.

The specific surface area of the exhon blacks about 30 to 1,000 m/g/m and the carbon black displays a distribution of both and 50 co 1,000 orange.

Sitting which our be employed are anarophous and have a primary peritied size of about 5 to 50 innomenteer and an except size of apparentation status (5 to 50 innomenteer and an except size of apparentation stall called size status 2 to about 120 micross. This alloss employed have a specific surface area of about 50 to 600 m²/gm and a dibut/phi/fluidatio (DEIP) discopring a beautiful source of the control of the control

Clays which can be employed ecoording to the Invention have an everage perticle size of about 0.01 to about 10 inflorons and a specific surface area of about 3 to 30 m2/gm. They exhibit oil absorption of about 20 to about 100 gms per 100 mms.

Organio polymeric substances wideh oan be used include polymera and copolymers of ethylene, propyleue, but an experience of ethics and polybytrone, in granuler or powder form. These organic polymeric materials fixed an everage particle star anging from about 0.01 to 100 microns, preferably 0.01 to 10 microns.

In general, the amount of fluidiscation and militared generally depends on the type of material utilized and the type of polybusidisms on the order to a facility about 15% to about 05%, and most professible about 15% to a about 45%, but and 15% to a about 15%, but and 15% to a about 15% but and 15% to a about 15% to a about

The Madication and can be introduced into the reador of or ament the top of the reador, at the bottom of the resetor, or to the recycle the decident the the bottom of the reador. Producely, the individual of all of introduced at or mise.

or the reador or subver the fidelized bed. It is preferred to year the fidelized read reight produced to read the produced to reador include and the produced to reador include an extension of the reador of the reador of the reador of the reador includes an include an extension of the reador of

A fluidized bed reaction system which is particularly sulled to production of polymeric materials in accordance with the present investion is illustrated in the drawing. With retarence thereto, the reactor 10 consists of a reaction zone 12 and evolutive reduction zone 14.

In general, the height to difference remains of the reaction zone can very in the range of about 2.7:1 to about 4.6:1. The range, of course, can vary to larger or smaller ratios and depends upon the desired production capacity. The cross-sectional area of the velocity reduction zone 14 is typically within the range of about 2.0 to about 2.0 multiplied by the cross-sectional area of the velocity reduction zone 14 is typically within the range of about 2.0 to about 2.0 multiplied by the cross-

sectional area of the reaction zone 12.

The reaction zono 12 Indixides a bed of growing polymer particles, brined pulymor particles and a minor amount of celetypa proteins indixided by the continuous files on polymerstable and modifying genous components in the form of metics up feed and recycle faild through the cention zono. To maintain a vielate fluidized bod, the supericleal gas verlocify through the bed modification proqued for fluidization, and professible as teles of 1.1 Macro solver minimum flow. Ordinarily, the superficial gas velocity does not exceed 5.0 fluidized and usually no more than 2.5 fluidized to the contraction of the contraction of

It is assential that the hest always contain particles to prevent the formulation of localized "hot spoic" and to enlarge and institute catalyst throughout the reaction zone. On feat up, the reaction is usually charged with a bad of particulate polymer perficts. Such particles may be federated in whate to be polymer for be formed or they may be different. When the other polymer for the time of the propriet of the fermion of the previous formed polymer periodes as the first product. Eventually, a fluitized bad of designed polymer periodes on the first product. Eventually, a fluitized bad of designed polymer periodes on the first product.

A partially or totally activated precursor composition and or catalyst used in the fluidized bod is preferably stored for sarvice in a reservoir 16 under a blanket of a gas which is inert to the stored material, such as rillrogen or argon.

Fluidization is achieved by a high rate of fluid recycle to and through the bed, hybrially on the order of about 50 the about 150 times the rate of feed of make-up fluid. The fluidized bed has the general appearance of a dense mass of inclividually moving particles as created by the percelation of gas through the bed. The pressure drop through file bad to equal to or slightly greater than the weight of the bad divided by the cross-sectional area. It is thus dependent on the account or of the means.

Make-up fluid can be fed to the bed at point 18. The composition of the make-up stream to determined by a gas analyzer 21. The gas analyzer characterises the composition of the recycle stream and the composition of the united by stream is extued a coordinately to maintain an essentially steady state assessor composition within the reaction zone.

The gas ambyzer is a conventional gas analyzer which operates in a conventional manner to determine the reception stream composition to facilities modificating the same description to facilities modificating the same composities. Such explainant to commercially a waitable from a wide writely of sources. The gas enelyzer 21 is tylorithy positioned to receive gas from a sampling point located between the windly recluding rough 1 and real accordance 24.8.

The higher bolling or readily condensable monomers can be introduced into the polymerization zone in various with both direction through a nozale (not shown in the dealing) but be bed or by repaining onto the top of the bed friency) in anozale (not otherwip positioned shown the bed, within any aid in eliminating come curryover of filese by the cycle paracterism. If fire rate of monomer feed is relatively small, header monomers can be infoduced into the polymerization) zone simply by successoration in the cycle use stream entering this bottom of the reason.

To ensure complete field called, the recycle stream and, where desired, part of the melve-up stream are returned incorph recycle in Fezz to the needs right part of the feet. There is pertendively agree desiriation printed 28 above the point of roturn to aid in fluidating the bot if, in passing through the bed, the recycle alream above the point of roturn to aid in fluidating the bot in passing through the bed, the recycle alream above the heat of reaction of sentrated by the portheristation for reaction.

The populs of the fluiditing element which has not reacted in the bed is removed from the polyametration cone, period why by possing it the to velocity requires one on the polyametration cone, period why by possing it the to velocity requires one of the polyametration and polyametration than bed. The necycle cheans is couperseased in a compressor 80 and their passed through a heat exchange or as the place is removed before it is returned to the bed. The heat exchange one has the placet in a tender and are also where the service are removed before it is returned to the bed. The heat exchange one has employed to lower the temperature of the origin search entering in stages. It is also possible to logical the compressor downstream from the heat exchanger or at intermediate point between several heat exchangem. After cooking, the recycle stream is returned to the reactor or at intermediate point between several heat exchangem. After cooking, the recycle stream is returned to the reactor it is the search or prevent contained polymer purified s from exiting out, augitorisating that on callel mass, and to prevent liquid in the cycle gas as distributed was the reactions between processes which contain liquid in the cycle gas as featured to the value of the contained at the contained at

The educated temperature of this best in maintained at an excentibly constant throughout or under attenty data continue by constant throughout the continue by constant throughout proceedings and the continued to the continued throughout throughout the continued throughout throughout throughout the continued throughout through

Cooliges distribution psays an important rote in the operation of the reactor. The flatitized basic various and formed are flocation polymer particides, as well no catalysis purioses. As the polymer particides are not and possibly active, they must be proceeded from celluling, for if a quiscount mass is allowed to cold, any active cellulyst continuous of the control of the contr

Gas distribution plate 28 is a preferred means for achieving good gas distribution and may be a screen, slotted plate, periorated plate, a plate of the bubble-cap type and the like. The elements of the plate may all be slationary, or

the pink may be of the mobile type disclosed in U.S. Patent No. 3,298,792. Whatever its design, it must diffuse the recycle fulld through the particles at the base of the best on keep the best in a fluidized condition, and also serve to support a guidecent best of real particles when the reactor is not in operation.

This preferred type of gas of strikulor plate 20 is motel and has holes distributed across its surface. The holes are normally of a dismoster of shout 1/2 into. The holes extend through the plate. Over each hole there is positioned a through a regular range to ten titualitied as 36 which is mounted on plate 28. The suggle hows serve to distribute the frow of high elong the curtace of the plate see as to world disquant zones of colids, in addition they prevent the polymer from flowing through the holes when the bud is suffice.

Any third hart to the cetalyst and reactions can also be present in the recycle stream. An activator compount, if utilized, is preliarably added to the reaction system downstream from heat exchanger 24, in which case the activator may be fed into the recycle system from dispenser 36 through line 40.

In the practice of this invention operating temperatures can extend over a range of from about -100°C to about 150°C with temperatures ranging from about 20°C to about 120°C being preferred.

The fluid bed reactor can be operated at pressures up to about 1000 psl and preferably at a pressure of from about 100 psl to about 600 psl. Operation at higher pressures favora heat transfer as an increase in pressure increases the unit volume heat capacity of the gas.

The partiality or foldily included precureor composition (e.g., neodywitten with an aligh halide) and/or catalyet (furginative collactive) ordered to an extensity is infected into the old at a public 24 which is above distinguisty plate 39 elevatives. He catalyst is injected at a point in the bed where good making with polymer particles occurs. Injecting the catalyst at a point above the distribution peter provides for existative or posterion of a fluitable debt polymerization resorts. Injecting the catalyst at a point above the distribution peter provides for existative or posterion or a fluitable debt polymerization resorts. Injection directly into the fluid state bed ricks in distributing the contribution of through the best and tends to exist the contribution within on consent for the form. Injection the catalyst little the resolar above the herican result in excessive adelyst carryover tool top recycle line where polymerization can occur in catalyst and one occur and or the fluid and these occurrence.

For a supported catalysi, if can be injected into the reactor by various techniques. It is preferred, however, to continuously feed the catalysis into the reactor officing a catalyse feedor as disclosed, e.g., in U.S. Patent No. 5,77,9712 For a catalyst in callotion, fliquid, or study from It is hypically individued as defeosed in U.S. Patent No. 5,317,9616 to Enady of all and U.S. Seriel No. 4,14,622, entitled "Process for Controlling Particle Growth during Production of Glictly Polymars," filed March 31, 1995. Both references are incorporated herein by reference. The catalysts is preferredly fell into the reactor at a point 20 o 40 personal of the reactor dismisler away from the reactor wall and at a height of about 5 to cloud 30 personal of this network of the hote.

A gais which is liner to the catalyst, such as rifrogen or argon, is preferably used to carry the catalyst into the bed. The rate of polymer production in the bud depende on the rate of catalyst injection and the concentration of morenec(s) in the reactor. The production rate is conveniently controlled by simply adjusting the rate of catalyst injection.

Since any change in the rate of catalysis injection will change the reaction rate and there the rate of which heat is generated in the best, the surpression of the recycle steem entering the reactor is equivaled upwards and observator to encommodate any change in the sate of heat generation. This ensures the mentiesence of an essentially constant to encommodate any change in the sate of heat generation. This ensures the mentiesence of an essentially constant is of course, useful to detect any temperature change in the bed on set to enable officer the operator or a conventional authomitatio control system to make a catalotic englishment in the temperature of the recycles determine.

Under in given net of oresetting conditions, the fluidized hed is maintained all essentiative countain height by withdrawing a partition of this bed up product at the ratio of formation of the particulate polymori product. Since the ratio of product formation, a measurement of the temperature rise of the fluid discress this research (the difference between their fluid intemperature) and exist influid remperature of the fluid discress and exist in the research of the difference between their fluid intemperature and exist influid remperature) is fortically so of the name of particular polymori formation in a constant influid velocity in our arresplicitle hospicative fluid is preparent in the intellitation.

On discharge of proficialise polymer product from resource 10, it is destant/who and prefeatable to exposure fluid from the product ent of bent in the recycle line 22. There are manners wave featant to the rate has complished this. One proferred systems is allow in it the demander. Thus, that and product leave reactor 10 at point 44 and order product discharge terms of the through wave 40, which may be a fell wave which the designed to have minimum subsidiors to the own or opported. Positioned above and below product discharge terms of a reactor variety and wave 50, 52 with the felter beling colleged to provide presenting of product into product any part to 14. The college terms of the product the product any part to 14. The college terms of product surge terms 65, i.e. a discharge variety of which within in the copre position identifies the conveying to shore, Valve 50 when 45 and 55 the complete that the complete that the conveying to shore. When 50 which within in the copre position identifies the product of conveying to shore, Valve 50 when 45 and 55 the conveying to shore, Valve 50 when 45 and 55 the conveying to shore, Valve 50 when 45 and 55 the conveying to shore and the conveying to shore of 45 and 55 these of the conveying to shore and the respective flow of the valve flow only a compressor 66 and their respective flow 25 through the 50 and the program of the 25 through the 50 and the program of the 25 through the 50 and the program of the 25 through the 50 and the program of the 25 through the 50 and the program of the 50 and 5

In a typical mode of operation, valve 48 is open and valves 50, 52 are in a closed position. Product and fluid enter product discharge lank 46. Valve 48 closes and the product is allowed to satisfa to product discharge tank 46. Valve 50 is then opened cernitish fluid to flow from product discharge tank 46 to surne tank 62 from which it is continually

present hack in or product in product in 22. Viries 50 is then closed and week 52 is operad an up product in product in Schwige product in 24. Viries 50. Viries 50 is then closed and week 16 is operad in upger with more than or product in product in 24. Viries 50 is then closed and week 16 is operad to increase and the product in 24. Viries 50 is in 45. Viries 50

The particular liming sequence of the valves is accomplished by the use of conventional programmable controllers which no well along in the art. Moreover, the valves can be kept abbitabilitiely free of agglomistated particles by directing a chosm of gas periodically through the valves and back for broader.

Another preferred product dischenge system which may be alternatively employed in the disclosion and chained in the oppending U.S. pacient application of Floor CE Anothernal Red July 28, 198, 198, 198, 198, 198 and earlief of Fluidized Bed Dischenge System's (now U.S. Palant No. 4, 292, 592). Such a system employe at least one (parallel) poir of lettak conording a selfation thick and a sunfer let less warranged in one non-hose high beginning the speaker seturated from the top of the suttling tank and a price thank warranged in one non-hose high beginning the speaker seturated from the top of the suttling tank to a point in the reactor near this top of the fluidized bod. Such alternative preferred product discharge system owleges the need for a recompression, time 64, 56, 68, as shown in the opsterned of the discharge.

The filedized bed reactor is coplained with an adequate vention system (not shown) to allow venting the bed during to chart up and dutil down. The reactor does not register the run of entiring entirer wast serging. The recycle late 22 end the elements therein (compressor 30, heat exchanger 24) should be amonth curricted and devoted of unnecessary obstitutions so a most to involve the flow of reactor fall that or entirely cutribles.

Illustrative of the polymers which can be produced in accordance with the invention are the following:

Polyisoprene

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Polystyrene Polybutadiene

SBR (polymer of butadiene copolymerized with styrene)

ABS (polymer of acrylonitrile butadiene and styrene)

Nitrile (polymer of buladiene copolymerized with acrylonitrile)

Butyl (polymer of isobutylane copolymerized with isoprene)

EPR (polymer of eitylane copolymentzed with propylene)

EPR (polymer of eitylane copolymentzed with propylene and a diene such as hexadiene, dicyclopentadiene, or subvidene notionnene).

Neoprene (polychloroprene)

Sificone (polydimethyl slickane)

Copolymer of othylone and vinyltrimethoxy allane

Copolymer of ethylene and one or more of acryonitrite, maleic acid esters, vinyl acetate, acrylic and methacrylic acid esters and the like

When it is clasted to produce polymers or copplymers using one or more monomers which ere all reliablyshy the poling or ready conductation and which term is given used to the the programmer and pressure conditions which are preferred for per phree fluiditized bud production in econograms with the investion, it is preferable to empty an intent advances with with remain gasepour useful are contained with the investion, and is preferable to enter the propose or intention and i

Nitrogen, because of the physical properties and relatively tow cost is a preferred medium for the mountacture of proymers from light prolling or reading condensation momens such as eigens, valy acides calls, approinting, manageylates, insilyjmenthocypiste and the like. Alkanes such as ethene and propone which remain gaseous at relatively for temperatures are also preferred.

Convolicinal techniques for the prevention of Souting of the reactor and polymer egiptomaristion can be used in the practice of our invention. Birustatives of these techniques or this interduction of flavely divided perituitation matter to provent agglomeration, as described in U.S. Patent Nos. 4,994,594 and 5,300,477; the addition of negative change generated to beliance positive voltages or the addition of positive change generated, changed changes to be added, either continuously or information to neutralized and production of the change generated promises to neutralized negative voltage portentials as described in U.S. Patent No. 4,803,251. Antichat substances may also be added, either continuously or informatificative for praving consistent of the present production.

The granular polyclustiene end/or polystomene elisationus of this trivertion can be compounded alone or in combination with other elisationers, e.g., antwart inther, styrane-butation rubbay, (halolpulyl midner, ethytane-propriendien antiber; reinfarcing tillers, e.g., authoritated, suffer; processing sids, entitlegraderist; and vulcantibing squinte using equipment and methods well busins to brow skilled in the art. It is hallowed that the such compounds, the singlety granuler form of the polybotaticine on polyboranes pengine laws or infantes in this wide the other electioner(s). But an would be

nehienable with correctional polytautetien or polytaporene to notice had form. It is generally desirable that distance blenks be inflantally mixed in order to optimize the mediantial projection of the vulcantzate. Furthermore, If the hear particulate magnitude this invention, used to maintain granutarity during and after the polymentration process, also happens to be a reinflording filler for the compound (e.g., candon black), then a further benefit may be actually reinflorded as therefore the project of the compound. This is because the filler, within normally would have of first the dragglomorated in the mixing process before it could be despersed, in this case orders the unitarity process streety substantially denoted memorated and disposition.

Elektronetic compounds pregared from granular polybutacleans and polytoprane or ministree through, are particularly useful as compounds of pregared from granular polybutacleans and polytoprane or ministree through a particular polybutaclean and polytoprane or ministree through a particular production of a radial usionability files, peopully formulated elektronetic compounds and his bed traded through in did to produce step stack for the tread, science, and bed filler components of the file, or to produce share stack for the unit retreating in maniferaction. Other specially formulated elektronetic compounds on the calmodronet of the side, or to produce cond-referred shall stack for the carcass and donunterestatib but components of this lot. The typeral or unusualized to ris but but the secretality file valores components of the side and tread or unusualized from the statistic or unusualized draw, andisky operanding and wiskly compressed the assembly to produce a todolic sleep, their place of the produce of the statistic or an explicit statistic produce, when the carbination of the statistic place of the statistic or the statistic place of the statistic place of the statistic place. The produce of the statistic place of the

When used as a constituent of the compounts, greater polyetusties of this invention particularly imports about no recisions, a faight exacting resistance, low here pleneration, and but coulding resistance. Greater polyetyperson of this invention particularly impairs building lock and green strength, which holitates he building and hardling of the green time, and term and cut resistance. The grautist, the relicionary polystactions and polysterian production that phase process of the present invention are also be omployed in other motified and extraded articles using techniques involve to these testile in the sur.

The following examples are provided to illustrate our invention.

Example 1

To a gas-phase siltred bed reactor that was maintained at a constant temperature of 60°C, 3.8 pounds of dried carbon black powder were added to act as a fluidization aid. To this was added 0.055 lbs TIBA, i.e. tritsobuly/aluminum. Then was ackled 1.86 lbs of 1.3-butadigns and sufficient nitrogen to bring the total reactor pressure to 315 psia. A small feed of supported catalyst consisting of neodymlum neodecancate on DEAC-treated silica was begun. Simultaneously, a small feed of 10 with triisobulylaluminum cocatalyst solution in isopentane was begun. Feed was adjusted to give a 7:1 molar ratio of Al:Nd. During e 2.8 hour polymerization reaction, a total of 6.93 lbs of additional butadiene were fed In order to replace butadiene that was polymerized or vented. A small vont stream leaving the reactor removed a total 49 of 0.95 lbs butadiene during the polymerization. At the end of the polymerization, the catalyst and co-catalyst feeds were stopped. The reactor was depressurized, and the reactor contents purged free of residual butadisna using ritrogen. The polymer was discharged from the reactor. The product did not contain any turnes that would indicate aggloriteration had occurred. To the contrary, the product was a free-flowing, fine, granular powder. The reactor was opened and deened to ensure that all product was recovered. The total weight of solid product that was recovered was adjusted for the carbon black that had been initially charged. The remainder (6.36 lbs) was the amount of butadiene polymer formed during the batch and which was present in the reactor when it was shut down. Since a total of 8 79 lbs (= 6.93 + 1.86) of builddiene were charged to the reactor and a total of 6.30 lbs (= 5.35 + 0.95) of butadiene have been accounted for leaving the reactor as polymer and in the continuous vent stream, there must have been 2.49 lise of butadiene monomer present to the reactor when polymerization was terminated. This monomer would have been removed 50 from the reactor when it was depressurized and the contents purged.

The reactor volume is GZ Filters (or 2.1 to July lene). All 60°C this vapor pressure of 1,6 that distinct is 10°C plant. The react of the future present in the reactor part age at a statistration would stop to 1,8 tillion. Of the total of 2.4 filt to set unpolyment to the total case of the react of 1,0 tillion was considered but indicated that was claim to perfect in the credit present in a condetence plants, for constructing, dissolved in this polyees mus. Thus, the reactor was being operated at a knaposature below the condensation temperature of the monomer present. The Cold Statist to Cliquid monomer combined with the SS is to or ophymer amounts of 1.1 the of condensed but-disner monomer per 100 his or (polyhindelme. Yet, the presence of little liquid monomer in the grey-place resector did not cause exponenciation of the polyhindelme.

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Properties of the above product are as follows:

Carbon black N-650 by analysis: 41%; Average particle size by sieve analysis: 0.025 inches; Neodyntium in polymer: 490 ppm; Reduced viscosity; 7.6 dfg; Cis 1-4: 97.6%

Examples 2-7 were conducted as in Example 1, but with the changes indicated in the tables.

Solidion Catalyst Preparation for Enempile 2. India a dry althogen purged Baskwas charged 12.39, grams of a hazana solution of recolymium readscarnost 6.5.44%. With hexane), To this was added 55 mL dry huxano. To this solution was added 3.0 mL of 1.5 M El₂ACI (1.0eq Alfrid). The mixture was stirred, charged to a prosunt/able metal cylindor and fed to the reactor as a solution.

Supported Catalyst Proporation for Example 5. To a 600 mL dry ritrogen purped fleek was added 78 15 gians of siles (notific activation) and 230 mL dry havens. Show, 40 mL of 1,5M EspAGI was added and to microre were alread for 40 minutes at room temperature. The solution was coded and 117 gians of a hexame codeling of programs of a feature of the solution of prodymbur versions (4,9 with 5 kil) was added slowly. The misture was alread for 30 minutes and then the solvent was removed under vectors.

	EXAMPLE NO.	2	3	4	5
6	PRODUCT:	POLYBUTADIENE	POLYBUTADIENE	POLYBUTADIENE	POLYBUTADIENE
	CATALYST DETAILS				
10	Catalysi	Neodymlum neo- desancele in hexeno	Neodynium versa- tate on DEAC-treated silica	Neodymium versa- tate on DEAC-treated silice	Neodymlum neo- decentrate on DEAG- treated silica
	Cocatelyst	10% TIBA in isopen- tene	10% TIBA in Isopen- tane	1:3 DIBAH: TIBA In Isopeniane	10% DIBAH in Iso- pentene
18	PROCESS CONDI- TIONS				
	Reaction Temp. (°C)	60	60	60	60
	Monomer part, pres- sure (psla)	63	63	63	63
20	Polymer produced (lb)	6.8	5.8 .	6.4	4.5
	Reaction time	5 hr	2 fir 30 min	2 hr 15 min	3 hr
25	PRODUCT ANALY- SIS				
	% Carbon Black N- 650 analysis	42	41	41	42
30	Avg. particlo size by sieve analysis (inch)	0 076	0.017	0.018	0.013
	Cocatalyst/Calalyst Feed ratio*	21	7	9.5	1 1
36	Nd content in poly- mer (ppm)	132	286	179	415
	Reduced Viscosity (dl/g)	12.8	103	7.6	4.9
40	Mooney viscosity (est. gum) ML (1 + 4 @ 100°C)				90
	% cls-1,4	99.1	97	96 2	97
			THE RESERVE OF THE PARTY OF THE		

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EXAMPLE NO.	6	7	ð
PRODUCT:	POLYBUTADIENE	POLYBUTADIENE	POLYISOPRENE
CATALYST DETAILS			
Caielyst 10	Neodymium neo- decancate on DEAG- treated silica	Naodymium neo- decancate on DEAC- treated silica	Neodymium neodecenoste on DEAC-treated allica
Countalyst	16% DIBAH in (copentane	10% DIBAH in Isopentane	10% TIBA in isopentane
PROCESS CONDITIONS			
Reaction Temperature (*C)	60	60	65
Monomer partial pressure (psla)	63	63	35
Polymer produced (lb)	5	4	
Reaction time	1 hr 45 min	1 hr 35 min	4
PRODUCT ANALYSIS		100000000000000000000000000000000000000	
% Carbon Black N-650 by analysis	36	39	40
Average particle size by sleve analysis (inch)	0.027	0.030	
Cocatelyst/Catelyst Feed ratio*	28	29	
Neodymium content in the polymer (ppm)	160	200	
Reduced Viscosity (dl/g)	4.2	9.7	
Mooney viscosity (est. gum)	62	39	
ML (1 + 4 @ 100°C) % cie- 1,4	96.5	95.6	

[&]quot; moler ratio of Al to rare earth motal in continuous feeds

Example 9

in an example of the process of the Invention a fluidized bed reaction system as described above, to operated as described below to process of the Invention and fluidized bed reaction system as described above, to operated as described below to proceed polyhoractions. The polymer is produced unrise the following reaction conditions of the reaction recording as the reaction of the process of the bedraids monomore involve the reaction is tip pair. The printing presence of all reliquinities of pairs and the process of the bedraids monomore involve the reaction system at the medical pairs and the process of the process of the bedraids are englisted to give a city in moder ratio of All of Mr. At all seally state to monomore is feel into the reaction system at seal of the process of the product that a Monomy viscosity Mr. (1 + 4/2) 100°Cy of 85. Other conditions are above for Example in into letter. At steady data into id Call, publication as the process of the process of the product that a Monomy viscosity Mr. (1 + 4/2) 100°Cy of 85. Other conditions are above for Example in this letter. At steady data into id Call, publications of the product that are above to the product that a Monomy viscosity Mr. (1 + 4/2) 100°Cy of 85. Other conditions are above for Example in this letter. At steady data into id Call, publication of 1.2 Mr. In the product that are also as the steady of the propriet of the first of the product of the product of the product that are also as the product of the product

The reactor volume is 55 fr³. At the partial pressure of 96 psia, there are 44.4 lbs of butadiene in the reactor gas-

phase. The leaf unpolymerized buildings in the reactor is that 85.0 the (a4.4. + 1.1.0), it all of this buildings were in the gate phase of this reactor at once it would have a partial pressure of 125 pala and its conclemation towardative would be 69°C. Therefore the reactor at 60°C is being operated below the condensation imprecative of the monomer present in the polymerization zone. Furthermore, the presence of this liquid monomer in the gas-phase reactor does not chause applicamentation of the colorest.

Example 10

In another example of the process of the invention the polymerization is conducted as described in Example 9 oxcept that the catalyst is needywitim needscendate fed as a solution in hexane. The table gives torther details on this example.

Example 11

In an exemple of the process of the Invention a Briddeach both reaction system an elecatibed above, is operated in described below to produce policysciene. The polymer is produced under the following reaction conditions 65°C concord rempenature and 100 pals total reactor prostore. The partial prosours of this isoprone monomore inside that reactor is 30 pals. The partial prosours of this isoprone monomore inside that reactor is 30 pals. The partial prosours of this isoprone monomore inside that reactor is 30 pals. The partial prosours of this isoprone monomore inside that reaction system of the consumption of this isoprone in this case is adjusted to give a 60°T moler ratio of 14 to 164. At steady state the monomore in feel into the reaction system at the ratio of 554 hbft, 1640 A 550 cathod beliefs feel for the reactor at the tast of 150 lbh. Exportan entormer lowers the reactor at 2 bits in vent streams. The production ratio is 30 lbh of polymers after adjusting for the carbon brack continuit. This product has a Monomy viscosity Min. 14. 46. 100°POS 65.00 lbh concretions necessitive of 15 mins of 150 lbh of polymers are shown to Example 11 in the table of 11 in the false.

At seady state a total of 56.4 in his payme is being fed to the reactor and a total of \$2 fair is excounted for leaving the reactor as gas in a vent diseam or as polymer. The officerance of 3.4 lbth must be unreacted liquid isopreme monomer in the polymer ferring the reactor. Since the polymer fed federaged is denicled with the polymer in the bed, the polymer for the bed must contain the samp proportion of liquid monomer, i.e. there must be 12.7 fine of dissolved liquid monomor in that 15 line pulsars bed.

The reactor volume is 6.5 ff². At the partial pressure of 80 pein, there are 17.2 to all incorrents in the reactor gazhues. The both unpolymerized isoperand in the reactor is flux 29.5 ft 6.7 (7.4 ± 1.2.7), it all of this isoprous were in the gas place of this reactor at one it would have a partial pressure of 54.5 pein and its condensation temperature would be 50°C. Threshort has reactor at 66°C is being operative below the condensation temperature of the monthment present in the polymetrization zone. Furthertenore, the presence of this liquid womener in the gas phase reactor does not consider condensation of the activate.

Example 12

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In another example of the process of the invention the polymerization is conducted as described in Example 11 except that the catalyst is neodymbon neodecanous fed as a solution in hoxane. The table gives further details on this example.

	EXAMPLE NO.	9	1 10	11	12
6	PRODUCT:	POLYBUTADIENE	POLYBUTADIENE	POLYISOPHENE	POLYISOPRENE
	REACTION CONDI- TIONS:				
10	Temperature (°C)	60	60	65	65
	Total Pressure (psla)	120	120	100	100
	Superficial Velocity (ft/s)	1.75	1.75	1.75	1.75
16	Production Rate (lb/h)	30	30	. 30	30
	Total Reactor Vol- ume (ft ⁵)	55	55	55	55
20	Reaction Zone Vol- ume (ff ³)	7.5	7.6	7.5	7.5
	Bed Height (ft)	7.0	7.0	7.0	7.0
	Bed Diameter (fi)	1.17	1.17	1.17	1.17
25	Red Weight (lbs)	112	112	112	112
	CYCLE GAS COM- POSITION (MOLE %):				
30	N ₂	20	20	70	70
	Buladiene	80	80		
	Isoprene			30	30
36	CATALYST:	Nd Neodecanoate on DEAC-treated sli- ica	Nd Neodecanoate in hexane	Nd Neodecanoste on DEAC-treated sil- ica	Nd Neodecanoste in hexerie
	CO-CATALYST:	TIBA	YIBA	TIBA	TIBA
40	Monomer Feed Rale (lb/h)				
	Butadiene	46,2	46.2		
	Isoprene			35.4	35.4
45	Mononier Vent Hate (lb/hr)	18	13	2	2
	POLYMEN COMPO- SITION (WL%);				
50	Buladiene	100	100		
	Isoprene			100	100

Clelms

65 1. A polymerisation catalyst comprising:

(A) rare earth metal component selected from the group consisting of (1) a reaction mixture obtainable by treating a affice support, a carbon black support or a mixed support thereof with a mono- or di-phyl aluminium obto-

- ride, adding a neodymlum carboxylate in the presence of a solvent to form a chury, and removing the solvent or (2) a feeding mixture obtainable by dissolving a neodymlum antipoxylate in a first solvent, adding silica or carbon black, and removing the solvent to obtain a dry coldr, contacting the dry solid with a solviet containing a none- or d-abyly abministure (chiotide in a second solvent, and removing the second solvent; and
- (B) a co-catalyst selected from the group consisting of a disobutyl aluminium hydride, trifsobutyl aluminium, and a mixture thereof.
 - A polymerication catalyst as claimed in claim 1 wherein the rare earth component (A) is obtainable by the sequential steps of:
 - (i) treating a silica support, carbon black support, or mixed support thereof with diathyl atominism chloride in a solvent to form a slurry;
 - (ii) adding a neodymiulm carboxylate to the slurry; and
 - (iii) removing the solvent.

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- A polymorisation catalyst as claimed in claim 1 wherein the rare earth component (A) is obtainable by the sequential steps of
- (i) Impregnating a silica support or a carbon black support with a neoxiymtum carboxylate in a first solvent to form a sturry;
 - (ii) removing the first solvent from step (i) to obtain a dry solid;
 - (iii) contacting sold dry solid with diethyl aluminium chloride in a second solvent to form a sharry; and
 - (iv) removing the second solvent from step (iii), wherein the first solvent of step (i) and the second solvent of step (iii) may be the same or different.
- A polymerisation catalyst as claimed in any one of the preceding claims wherein the neodyntum carboxylate is neodyntum neodecanoate, neodyntum octosta or neodyntum versatiste.
- A polymerication catalyst as claimed in any one of the preceding claims wherein the organic alityl aluminium compound is distinyl aluminium chloride.
- A polymerisation catalyst as claimed in claim 2 wherein the solvent is n-hexene or cyclohexene.
- A polymerisation catalyst as claimed in claim 3 wherein the first solvent is n-hexane or cyclohexane and the second colvent is a hydroparbon.

